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SURFACE PLASMON RESONANCE SENSOR

This invention relates to a Surface Plasmon Resonance
Sensor. In particular it relates to an improved design

5 of Surface Plasmon Resonance Sensor that is compact,

6 simple to align and cost effective to produce, thus

making it ideal for field applications.

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The phenomenon of Surface Plasmon Resonance (SPR) is well known to those skilled in the art having being first demonstrated over twenty five years ago. Surface Plasmon Resonance is a charge-density oscillation that may exist at the interface of two media that exhibit dielectric constants of opposite signs, for example a metal and a

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dielectric.

17 Surface Plasmon Resonance sensors described in the Prior art generally comprise an optical system, a transducing 18 medium that generally combines the optical system and the 19 relevant chemical or biochemical domains, and an 20 21 electronic system that supports the optoelectronic components of the sensor, and allows for the required data 22 23 processing. The devices come in three main 24 configurations namely:

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1 Prism coupler based systems; (1)

becomes incident on a detector 7.

- 2 (2) Grating coupler based systems; or
- 3 Optical waveguide based systems. (3)

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5 A typical prism coupler based system 1 is presented 6 schematically in Figure 1. This system is generally 7 accepted as being the best suited for sensing and 8 therefore has become the most widely employed system in 9 the art. In this configuration a light wave 2 passes 10 through a first element of an optical system 3 before 11 passing into a prism 4. Thereafter, the light wave 2 12 experiences total internal reflection at the interface 13 between the prism 4 and a thin metal layer 5 (typically 14 of a thickness of around 50 nm). The light wave 2 then 15 passes through a second element of the optical system 6 16

that acts to manipulate the light wave 2 such that it

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19 The Surface Plasmon Resonance sensor 1 is an ideal medium 20 for analysing samples that become attached to the metal 21 layer 5. SPR is a phenomenon that occurs when light 22 incident upon the metallic layer 5 provides an absorption 23 energy capable of vibrationally exciting the packets of 24 electrons (or plasmons) located on the surface of the metal layer 5. As such the energy required to achieve 25 26 SPR is highly dependent upon the dielectric constant of 27 the species at the surface of the metal, the wavelength 28 of the light wave 2 and the angle of incidence of the 29 light wave 2.

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31 is known in the art the use of a particular 32 monochromatic light source of a known wavelength incident 33 at variable angles, or across a range of known angles, 34 allows a reference Reflectance Angle versus Intensity

l data to be recorded. The presence of any foreign bodies

- 2 that become attached to the surface of the metal layer 5
- 3 then act to change the value of the dielectric constant
- 4 experienced by the light wave 2 at the surface of the
- 5 metal layer 5. As such the presence of these foreign
- 6 bodies can be easily detected and thereafter quantified
- 7 by monitoring the profile of the Reflectance Angle versus
- 8 Intensity curves.

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- 10 The systems described in the Prior Art are difficult to
- 11 optically align and so require a skilled operator.
- 12 Furthermore the systems are not easily miniaturised and
- 13 as such are not easily adapted to be used as field based
- 14 instruments. Generally, a user is required to take a
- 15 sample that then needs to be taken to the laboratory for
- 16 testing by the operator. This process can lead to
- 17 significant delays in obtaining results. Such delays can
- 18 be fatal when the instrument is employed as a biosensor
- 19 to detect particular pathogens.

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- 21 It is an object of an aspect of the present invention to
- 22 provide a Surface Plasmon Resonance Sensor that overcomes
- 23 one or more of the limiting features associated with the
- 24 apparatus and methods described in the prior art.

- 26 According to a first aspect of the present invention
- 27 there is provided a cartridge for use in a Surface
- 28 Plasmon Resonance sensor, the cartridge comprising an
- 29 optical element having a first surface and a mounting
- 30 member for supporting a sensing agent located on a second
- 31 surface of the optical element wherein the first surface
- 32 comprises a first means for directing a beam of light
- 33 incident on the optical element towards the second
- 34 surface at an angle of incidence to the second surface

that results in substantially total internal reflection

- 2 of the beam of light at an interface of the mounting
- 3 member and the second surface.

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- 5. Most preferably the optical element further comprises a
- 6 third surface for the exit of the beam of light from the
- 7 optical element wherein the third surface includes a
- 8 second means for directing the beam of light.

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- 10 Preferably the optical element comprises a material
- 11 having a first dielectric constant while the mounting
- 12 member comprises a material having a second dielectric
- 13 constant wherein the second dielectric constant is of an
- 14 opposite sign to that of the first dielectric constant.

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- 16 Most preferably the first means for directing the light
- 17 beam comprises a focusing element for focusing the beam
- 18 of light to a line at the interface of the mounting
- 19 member and the second surface.

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- 21 Preferably the second means for directing the light beam
- 22 comprises a defocusing element.

23

24 Preferably the mounting member comprises a metal.

25

- 26 Preferably the optical element comprises an injection
- 27 moulded plastic material.

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- 29 Most preferably the sensing element comprises one or more
- 30 antibodies each antibody being suitable for binding a
- 31 pathogen.

- 33 Preferably the bound pathogen is selected from the group
- 34 comprising Legionella, Escherichia coli, Salmonella,

l Bacillus Anthracis, Yersinia Pestis, Lysteria,

- 2 Cryptosporidium, Variola virus, Picomaviridae Apthovirus,
- 3 Filoviruses, any plasticiser, steroid, medicinal drug or
- 4 illicit substance or any other known fluid borne
- 5 bacterium.

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- 7 Preferably a protein substrate and a ligand is employed
- 8 to bind a biotinylated antibody to the metal.

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10 Preferably the protein substrate comprises biotin.

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- 12 Preferably the ligand comprises a protein selected from
- 13 the group comprising avidin, strepavidin and neutravidin.

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- 15 According to a second aspect of the present invention
- 16 there is provided a Surface Plasmon Resonance sensor
- 17 comprising a light source for generating a beam of light,
- 18 a cartridge according to the first aspect of the present
- 19 invention, a channel suitable for containing a fluid
- 20 sample to be tested and a light beam detection means
- 21 wherein the employment of the cartridge allows for the
- 22 miniaturisation of the sensor.

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24 Most preferably the light source comprises a diode laser.

25

- 26 Preferably the channel locates on the second surface of
- 27 the cartridge such that the fluid sample contained within
- 28 the cartridge makes physical contact with the mounting
- 29 member.

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- 31 Preferably the light beam detection means comprises a
- 32 detector and a data processing means.

1	According to a third aspect of the present invention
2	there is provided a method of field detection of one or
3	more pathogens comprising the steps of:
4	1) Selecting an appropriate cartridge for the
5	detection of one or more pathogens for use in a
6	Surface Plasmon Resonance sensor;
7	2) Calibrating the Surface Plasmon Resonance sensor;
8	and
9	3) Testing a fluid sample for the presence of one or
0	more of the pathogens;
1	
2	Preferably the selection of the appropriate cartridge
13	comprises locating the cartridge with one or more
4	appropriate antibodies for binding with the one or more
15	pathogens.
16	
17	Preferably calibrating the Surface Plasmon Resonance
8	sensor comprises:
19	1) Irradiating the mounting member with the beam of
20	light in the absence of the fluid sample; and
21	2) Detecting a component of the beam of light
22	reflected from the mounting member and storing the
23	data as a reference signal;
24	•
25	Preferably testing of a fluid sample for the presence of
26	one or more pathogens comprises:
27	1) Locating the fluid sample with respect to a
28	channel;
29	2) Connecting the channel to the cartridge;
30	3) Irradiating the fluid sample with the beam of
31	light;
32	4) Detecting the beam of light reflected from the
33	mounting member and storing the data as a sample
34	signal; and

5) Comparing the sample signal with the reference 2 signal. 3 Embodiments of the invention will now be described, by way of example only, with reference to the accompanying 5 drawings, in which: 8 Figure 1 present a prism coupler based Surface 9 Plasmon Resonance sensor as described in 10 the Prior Art; 11 Figure 2 present a disposable cartridge 12 Surface Plasmon Resonance sensor 13 accordance with an aspect of the present 14 invention; 15 Figure 3 present a schematic representation of the 16 Surface Plasmon Resonance sensor of 17 Figure 2; and 18 Figure 4 present a schematic representation of a 19 binding method employed by the Surface 20 Plasmon Resonance sensor of Figure 2; and 21 Figure 5 presents typical Angle versus Intensity 22 curves as may be obtained by the Surface 23 Plasmon Resonance sensor. 24 Figures 2 and 3 present a disposable cartridge based 25 Surface Plasmon Resonance sensor 8 in accordance with an 26 aspect of the present invention. The sensor can be seen 27 to comprise a diode laser 9, a disposable cartridge 10 28 and a charge coupled device (CCD) detector 11 that is 29 **30** connected to a data processing unit 12. 31

The disposable cartridge 10 comprises a shaped entrance 32 surface 13, a shaped exit surface 14 and a gold strip 15 33

that is attached to a third side of the disposable 34

1 cartridge 16. A channel 17 is employed to enclose the

- 2 gold strip so providing a means for containing and
- 3 introducing a fluid sample to the surface of the gold
- 4 strip 15. The disposable cartridge 10 can be detached
- 5 from the channel 17 so as to enable the cartridge 10 to
- 6 be disposed of and replaced, as required.

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- 8 In order that the cartridge 10 be correctly aligned to
- 9 the diode laser 9, the CCD detector 11 and located
- 10 correctly with the channel 17, the channel 17 may further
- 11 comprise either male of female members (not shown) that
- 12 interact with female or male members, respectively,
- 13 located on the surface of the cartridge 10.

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- 15 For the Surface Plasmon Resonance sensor 8 to operate
- 16 correctly there must be a means whereby the relevant
- 17 pathogen 18 to be detected can attach to surface of the
- 18 gold strip 15. There are several techniques known to
- 19 those skilled in the art for binding pathogens 18 to a
- 20 metal strip.

- 22 Figure 4 presents a schematic representation of a binding
- 23 method suitable for use with the Surface Plasmon
- 24 Resonance sensor 8. The first stage involves binding a
- 25 suitable protein substrate 19, for example biotin, to the
- 26 surface of the gold strip 15. Stage two involves
- 27 attaching a ligand 20 to the protein substrate 19. A
- 28 suitable ligand 20 for conjugating with biotin is avidin
- 29 although steptavidin or neutravidin may also be employed.
- 30 The third stage then involves the attachment of an
- 31 antibody 21, appropriate for the relevant pathogen 18 to
- 32 be tested for, to the ligand 20. This attachment is
- 33 achieved by employing antibodies 21 that have been
- 34 biotinylated 22.

When the gold strip 15 has been treated as described above the Surface Plasmon Resonance sensor 8 is ready for 3 use. The diode laser 9 provides the required light beam 5 The light beam 23 is focused to a line 24 on the 23. gold strip 15 on passing through the shaped entrance surface 13. This provides a large area of interaction between the light beam 23 and the gold strip 15. Such an area of interaction allows a range of spatially resolved biotinylated antibodies 22 to be deposited on a single 10 11 cartridge 10. The light beam 23 is then totally internally reflected so as to traverse through the shaped 12 exit surface 14. This results in the light beam 23 being 13 defocused such that the incident signal from each of the 14 biotinylated antibodies 22 is spatially resolved across 15 the whole area of the CCD detector 11. Data processing 16 17 then carried out on the detected signal, as is 18 appropriate.

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Figure 5 presents a schematic Reflectance Angle versus 20 Intensity curves typically obtained by the Surface 21 22 Plasmon Resonance sensor 8. The solid curve 25 corresponds to the case where no pathogen 18 is present 23 24 in the fluid sample as indicated in Figure 5(a). However, Figure 5(b) shows the case when a pathogen 18 is 25 present in the fluid sample, as represented by the broken 26 27 curve 26. The pathogen 18 on becoming attached to the surface of the gold strip 15 alters the value of the 28 dielectric constant experienced by the light beam 23 at. 29 the surface of the gold strip 15. As such the presence 30 of the pathogen 18 alters the profile of the Angle versus 31 Intensity curve, so permitting quick and easy detection 32 of the presence of the pathogen 18. 33 34

The employment of the disposable cartridge 10 and a diode

laser 9 light source provides the Surface Plasman

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2 laser 9 light source provides the Surface Plasmon
3 Resonance sensor 8 with significant inherent advantages

4 over those taught in the Prior Art. In the first

5 instance these elements significantly simplify the

6 optical alignment requirements of the device as well as

7 allowing for the significant miniaturisation of the

8 device. As such, the Surface Plasmon Resonance sensor 8

9 provides a compact, simple to align and cost effective

10 device for the field testing of the presence of a

11 pathogen 18. The miniaturisation of the device has the

· 12 added advantage that it increases the sensitivity of the

13 sensor since all of the functionalised area of the gold

14 strip 15 can be contained within the focused line 24 area

15 of the incident light beam 23.

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17 In particular, the fact that the focusing and defocusing 18 elements are incorporated directly within the disposable 19 cartridge 10 simplifies the time consuming alignment 20 requirements associated with the optical systems 3 and 6 21 of the Prior Art sensors. In addition, the employment of 22 an injection moulding technique allows for the low cost fabrication of the disposable cartridge 10. 23 24 technique therefore makes it cost effective to remove and 25 dispose of the cartridge 10 after use and simply replace 26 it with a new cartridge 10, as required. The use of 27 these disposable cartridges 10 significantly reduces the 28 time consuming cleaning requirements associated with the 29 sensors described in the Prior Art.

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An alternative embodiment of the Surface Plasmon Resonance sensor (not shown) the fluid sample to be tested is continuously passed through the channel 17 and across the surface of the gold strip 15. This allows for

the Surface Plasmon Resonance sensor to continuously

2 monitor a fluid source for the presence of a pathogen 18

rather than testing a single sample taken from the fluid

source as discussed in relation to the above preferred

5 embodiment.

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7 The Surface Plasmon Resonance sensor 8 described herein

8 is particularly suitable for the detection of the

bacteria Legionella in water samples obtained from

10 industrial or recreational sources. This is of

11 particular importance in evaluating and controlling the

12 risk to public health presented by the often-fatal

13 condition Legionnaires disease and the less serious but

14 far more common condition of Pontiac Fever. Existing

15 techniques are either very slow or too labour intensive

16 to meet market demands, since they generally require

17 qualified microbiologists to perform testing at

18 specialist laboratories.

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20 The availability of the focused line 24 interaction area

21 on the gold strip 15 allows for the functionalisation of

22 the interaction area for different antibodies that are

23 sensitive to different forms of the Legionella bacteria.

24 Thus, the above apparatus provides a sensor that is

25 capable of simultaneously detecting and discriminating

26 between Legionella pnuemophilla serogroup 1 and

27 Legionella serogroups 2-15.

28

29 Although ideal for the detection of the bacteria

30 Legionella, it will be obvious to one skilled in the art

31 that the surface Plasmon Resonance sensor may be easily

32 adapted for use in the detection of alternative species

33 e.g. Escherichia Coli, Salmonella, Bacillus Anthracis,

34 Yersinia Pestis, Lysteria, Cryptosporidium, Variola

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l virus, Picomaviridae Apthovirus, Filoviruses, any

2 plasticiser, steroid, medicinal drug or illicit substance

3 or any other known fluid borne pathogen.

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5 In addition to the use for water quality monitoring as

6 described above it would be obvious to one skilled in the

7 art that the Surface Plasmon Resonance sensor 8 is also

ideal for use in healthcare, especially for use as a

9 point of care diagnostic.

10

11 Aspects of the present invention described above offer

12 significant advantages over the Prior Art. In the first

13 instance the Surface Plasmon Resonance sensor provides a

14 compact, simple to align and cost effective device for

15 the field testing of the presence of a pathogen. The

16 device is ideal for the expeditious detection and

17 identification of a range of pathogens. Further, the

18 incorporation of the focused line area provides a means

19 for carrying out such a detection and identification

20 process simultaneously for a number of different

21 pathogens.

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23 The foregoing description of the invention has been

24 presented for purposes of illustration and description

25 and is not intended to be exhaustive or to limit the

26 invention to the precise form disclosed. The described

27 embodiments were chosen and described in order to best

28 explain the principles of the invention and its practical

29 application to thereby enable others skilled in the art

30 to best utilise the invention in various embodiments and

31 with various modifications as are suited to the

32 particular use contemplated. Therefore, further

33 modifications or improvements may be incorporated without

1 departing from the scope of the invention herein

2 intended.